## **BLAST Basic Local Alignment Search Tool**

Your search parameters were adjusted to search for a short input sequence.

Edit and Resubmit Save Search Strategies. Formatting options Download

Blast 2 sequences

#### SEQ ID NO: 3

Results for: Icl|11229 None(21bp)

Your BLAST job specified more than one input sequence. This box lets you choose which input sequence to show BLAST results for.

Query ID

|c||11229

Description
None

Molecule type
nucleic acid

Query Length 21

Subject ID

gi|1914699|emb|X98077.1| Description

Hepatitis B virus complete genome, wild type

Molecule type nucleic acid

Subject Length

Program BLASTN 2.2.22+ Citation

#### Satorano

Stephen F. Altschul, Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402.

Other reports: Search Summary [Taxonomy reports]

## **Search Parameters**

Program	blastn
Word size	7
Expect value	1000
Hitlist size	100
Match/Mismatch scores	1,-3
Gapcosts	5,2
Filter string	F
Genetic Code	1

## Karlin-Altschul statistics

Params	Ungapped	Gapped
Lambda	1.37406	1.37406
K	0.710603	0.710603
Н	1.30725	1.30725

# **Results Statistics**

Effective search space 44912

Designing or Testing PCR Primers? Try your s

Graphic Summary

## Distribution of 8 Blast Hits on the Query Sequence

[?]

An overview of the database sequences aligned to the query sequence is shown. The score of each alignment is indicated by one of five different colors, which divides the range of scores into five groups. Multiple alignments on the same database sequence are connected by a striped line. Mousing over a hit sequence causes the definition and score to be shown in the window at the top, clicking on a hit sequence takes the user to the associated alignments. New: This graphic is an overview of database sequences aligned to the query sequence. Alignments are color-oded by score, within one of five score ranges. Multiple alignments on the same database sequence are connected by a dashed line. Mousing over an alignment shows the alignment definition and score in the box at the top. Clicking an alignment displays the alignment detail.

		Color key for	alignment sc	ores	
<40 Query	40	-50	500.00	90-200	>=200
0	4	8	12	16	20

100% 9e-09 100%



## Plot of Icl|11229 vs gi|1914699|emb|X98077.1| [?]

This dot matrix view shows regions of similarity based upon the BLAST results. The query sequence is represented on the X-axis and the numbers represent the bases/residues of the query. The subject is represented on the Y-axis and again the numbers represent the bases/residues of the subject. Alignments are shown in the plot as lines us trand and protein matches are slanted from the bottom left to the upper right corner, minus strand matches are slanted from the upper left to the lower right. The number of lines shown in the plot is the same as the number of alignments found by BLAST.

×

Descriptions

Legend for links to other resources: UniGene GEO Gene Structure Map Viewer

Sequences producing significant alignments:

(Click headers to sort columns)

X98077.1 Hepatitis B virus complete genome, wild type 42.1 154

Sort alignments for this subject seq E value Score Percent identity Query start position Subject star

```
Designing or Testing PCR Primers? Try your's
```

Alignments Select All Get selected sequences Distance tree of results Multiple alignment. NEW

```
>emb|X98077.1| Hepatitis B virus complete genome, wild type
Length=3215
 Score = 42.1 bits (21), Expect = 9e-09 Identities = 21/21 (100%), Gaps = 0/21 (0%)
 Strand=Plus/Minus
               GTGCAGAGGTGAAGCGAAGTG 21
Query 1
Sbjct 1602 GTGCAGAGGTGAAGCGAAGTG 1582
 Score = 22.3 bits (11), Expect = 0.009 Identities = 11/11 (100%), Gaps = 0/11 (0%)
 Strand=Plus/Minus
               GCAGAGGTGAA 13
Query 3
Sbjct 1834 GCAGAGGTGAA 1824
 Score = 16.4 bits (8), Expect = 0.54 Identities = 8/8 (100\%), Gaps = 0/8 (0\%)
 Strand=Plus/Minus
Query 2 TGCAGAGG 9
Sbjct 410 TGCAGAGG 403
 Score = 16.4 bits (8), Expect = 0.54
 Identities = 8/8 (100%), Gaps = 0/8 (0%)
 Strand=Plus/Minus
Query 6
               GAGGTGAA 13
Sbjct 2051 GAGGTGAA 2044
 Score = 14.4 bits (7), Expect = 2.1 Identities = 7/7 (100\%), Gaps = 0/7 (0\%)
 Strand=Plus/Minus
Query 3
               GCAGAGG 9
Sbjct 1261 GCAGAGG 1255
 Score = 14.4 bits (7), Expect = 2.1
 Identities = 7/7 (100%), Gaps = 0/7 (0%)
 Strand=Plus/Minus
Query 5
               AGAGGTG 11
Sbjct 1533 AGAGGTG 1527
 Score = 14.4 bits (7), Expect = 2.1 Identities = 7/7 (100%), Gaps = 0/7 (0%)
```

Select All Get selected sequences Distance tree of results Multiple alignment NEW